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(54) Title: UNSATURATED α-OLEFIN COPOLYM	IERS A	AND METHOD FOR PREPARATION THEREOF
(57) Abstract		
erally crystalline, free of gel and cross-links, and conta	in unsa nd may	ers, and a method of preparation thereof. The copolymers are genturated side chains and long chain branching. The copolymers conbe prepared by copolymerization using a solid-phase, insoluble cone melting point of the copolymer.

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# UNSATURATED $\alpha$ -OLEFIN COPOLYMERS AND METHOD FOR PREPARATION THEREOF

## Field of the Invention

This invention relates to copolymers of  $\alpha$ -olefins and  $\alpha, \omega$ -dienes, and a method for their preparation. More particularly, the invention relates to crystalline, gel-free, long chain branched and unsaturated  $\alpha$ -olefin/ $\alpha, \omega$ -diene copolymers and their preparation.

#### Relevant Art

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Selected dienes have been copolymerized with aolefins in order to introduce unsaturation into the 10 polymer as a site for chemical reactions such as crosslinking. In U. S. Patent 3,658,770 to Longi et al, it is taught that these dienes should be non-conjugated and should have two different kinds of double bonds, the first double bond being copolymerizable with the  $\alpha$ -15 olefin and the second double bond being noncopolymerizable by Ziegler-type catalysts. Thus, useful classes of dienes for introduction unsaturation into ethylene-propylene terpolymers are straight-chain and branched-chain 20 acyclic single ring alicyclic dienes, and multi-ring alicyclic and bridged ring dienes, wherein one of the double bonds will copolymerize while the second is inert to the catalyst. Such useful dienes are 1,4-hexadiene, 25 dicyclopentadiene and 5-ethylidene-2-norbornene.

The copolymerization of branched 1,4-dienes with ethylene, propylene or 4-methyl-1-pentene is similarly described in U. S. Patent 4,366,296 to Kitigawa et al. While the copolymerization of propylene with α,ω-dienes of at least 6 carbon atoms using a Ziegler type catalyst in a hydrocarbon diluent is taught in U. S. Patent 3,351,621 to Bacskai, the copolymerization has invariably led to a copolymer insoluble in hot

decahydronaphthalene, indicative of the presence of cross-links and gel.

U. S. Patent 2,933,480 to Gresham et al, discloses the copolymerization of  $\alpha$ -olefins and diolefins in which both double bonds are terminal to obtain an elastomeric (non-crystalline) interpolymer. From U. S. Patents 4,551,503; 4,340,705; and 3,991,262, all to Lal et al, it is known to prepare interpolymers of  $C_4$ - $C_{12}$   $\alpha$ -olefins with  $C_8$ - $C_{36}$   $\alpha$ , $\omega$ -dienes using a Ziegler-type catalyst specially treated with hexaalkyl phosphoric triamides or organophosphate esters. However, the interpolymers taught in these patents are also elastomers, and not crystalline thermoplastics.

U. S. Patent 3,480,599 to Park discloses 15 elastomers made by copolymerizing trienes with  $\alpha$ -olefins.

As far as applicants are aware, there is no prior art teaching crystalline copolymers of  $\alpha$ -olefins such as propylene with  $\alpha, \omega$ -diene, wherein the copolymer is essentially gel-free and contains residual reactive unsaturation. Nor are applicants aware of any prior art teaching the copolymerization of  $\alpha$ -olefins and  $\alpha, \omega$ -dienes to produce a crystalline, gel-free copolymer, wherein the copolymerization is effected without a diluent, below the glass transition temperature of the copolymer product, and using a solid heterogeneous catalyst.

## Summary of the Invention

It has been discovered that dienes having two Ziegler-polymerizable double bonds can be copolymerized with α-olefins to obtain a crystalline structure without extensive gel formation if the copolymerization is effected with a Ziegler-type catalyst having an immobile or insoluble catalyst site under conditions for rapidly immobilizing the polymer chain. Immobile catalyst sites are, for example, those on the face of

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crystals, or on surfaces of solid-supported catalyst. The growing polymer chains may be immobilized by crystallization and/or by copolymerizing near or below the glass transition temperature of the polymer.

In one aspect, the invention provides a method of preparing a gel-free, unsaturated copolymer of an αolefin and an  $\alpha, \omega$ -diene. The method comprises charging to a reaction zone  $\alpha$ -olefin having from 2 to about 12 carbon atoms,  $\alpha, \omega$ -diene having from 7 to about 30 10 carbon atoms, and solid heterogeneous coordination catalyst. The method also includes maintaining the reaction zone essentially free of solvent and at a temperature at which the a-olefin is fluid but below crystalline melting point of the copolymer, effective to produce an essentially gel-free copolymer with unsaturation.

In another aspect, the invention provides an unsaturated crystalline a-olefin copolymer. The copolymer comprises  $\alpha$ -olefin interpolymerized with from 20 0.01 to 5 mole percent of an  $\alpha, \omega$ -diene having from 7 to about 30 carbon atoms to produce an unsaturated crystalline copolymer essentially free of gel. copolymer of this invention incorporates the  $\alpha, \omega$ -diene comonomer in two different ways. First, one end of the 25 diene comonomer may be reacted in the α-olefin backbone leaving the other end unreacted to provide unsaturated Second, the other end of the diene side chains. comonomer may react with additional monomer to provide long side chain branches greater in length than the 30  $\alpha, \omega$ -diene, in a fashion analogous to the branching in low density polyethylene (LDPE) which is also sometimes referred to as branched polyethylene or high pressure polyethylene. The predomination of either manner of diene comonomer incorporation can be effected by the 35 selection of catalyst and polymerization conditions, and the copolymer properties tailored accordingly. side chain unsaturation, for example, imparts gamma

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radiation stability and reactive functional sites for chemical modification, whereas long chain branching imparts greater melt strength and elasticity to facilitate processing the copolymer in heretofore difficult polypropylene applications such as, for example, blow molding, stretch blow molding, extrusion into pipe, extrusion coating, and high temperature flexural modulus retention. Surprisingly, however, the  $\alpha$ -olefin- $\alpha$ ,  $\omega$ -diene copolymer is free of xyleneinsoluble gel, regardless of which type of diene incorporation predominates.

### Detailed Description of the Invention

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The present invention involves the reaction of an  $\alpha$ -olefin with an  $\alpha, \omega$ -diene. Suitable \alpha-olefins include, for example, ethylene, propylene, butene-1, 15 pentene-1, hexene-1, heptene-1, 4-methyl-1-pentene, 4methyl-1-hexene, 5-methyl-1-hexene, 1-octene, 1-decene, 1-undecene, 1-dodecene and the like. ethylene, propylene, butene-1, pentene-1, and 4-methyl-**2**C 1-pentene are preferred, and propylene is particularly In addition, mixtures of these  $\alpha$ -olefins which result in crystalline polymers may also be used, such as, for example, propylene and a minor proportion of ethylene. The invention is described below with reference to propylene as an example of the  $\alpha$ -olefin sake of brevity and clarity with the understanding that other α-olefins are also contemplated.

Suitable  $\alpha, \omega$ -dienes contain at least 7 carbon atoms and have up to about 30 carbon atoms, with from 8 to 12 carbon atoms being preferred. Representative examples of such  $\alpha, \omega$ -dienes include 1,6-heptadiene, 1,7-octadiene, 1,8-nonadiene, 1,9-decadiene, 1,10-undecadiene, 1,11-dodecadiene, 1,12-tridecadiene, 1,13-tetradecadiene, and the like. Of these, 1,7-octadiene,

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and 1,9-decadiene are preferred, particularly 1,9-decadiene.

The a,w-diene may comprise from about 0.01 to about 5 mole percent of the copolymer, preferably from 5 about 0.05 to about 2 mole percent, and especially from about 0.1 to 1 mole percent. The diene content can be estimated, for example, by measuring absorbance at 722 cm<sup>-1</sup> using infrared spectroscopy. Depending upon the catalyst, polymerization conditions and comonomer selection, the copolymer may incorporate unsaturated comonomer side chains comprising from about 0.001 to about 5 mole percent of the copolymer, preferably from about 0.01 to about 2 mole percent, and especially from about 0.1 to about 1 mole percent.

15 sharp contrast to the prior art diene however, the present copolymers, copolymer is essentially gel-free and generally crystalline. absence of gel and cross-links in the present copolymer is evidenced by, for example, very low (less than 0.1 20 weight percent) or undetectable xylene-insoluble fractions. Also, the high crystallinity of the present copolymers is evidenced by a relatively high hot nheptane insolubles (HI) fraction, preferably at least 80 weight percent up to as much as 95 weight percent or 25 more. Further, the present copolymers are branched and may have a branching index from about 0.2 to about 0.9, preferably from 0.6 to 0.8. As used herein, the branching index is defined as the ratio of intrinsic viscosity of the  $\alpha, \omega$ -diene- $\alpha$ -olefin copolymer 30 to the intrinsic viscosity of the poly-α-olefin homopolymer having same molecular the distribution as the copolymer. The copolymer may have any molecular weight suitable for its intended end use, and in the case of polypropylene copolymers, preferably 35 has a melt flow rate (MFR) of from about 0.01 to about 2000, more preferably from about 0.1 to about 300 and especially from about 0.5 to about 70. As used herein,

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MFR is determined according to ASTM D-1238, condition L (2.16 kg, 230°C).

 $\alpha, \omega$ -Diene- $\alpha$ -olefin copolymers are prepared according to the method of the invention 5 interpolymerizing the α-olefin and the diene under conditions in which the catalyst site relatively insoluble and/or immobile so that the polymer chains are rapidly immobilized following their formation. Cross-link formation through 10 unsaturation remaining at the free-end of the branch chain resulting from incorporation of the diene into the copolymer is thus inhibited by rapidly immobilizing the otherwise reactive unsaturation site. immobilization is effected, for example, by (1) using a 15 solid, insoluble catalyst, (2) conducting copolymerization in a medium in which the resulting copolymer is generally insoluble, and (3) maintaining the polymerization reactants and products below the crystalline melting point of the copolymer.

Catalysts suitable in the method of the present 20 invention include solid, heterogeneous coordination catalysts, such as Ziegler-type catalysts. insoluble or otherwise solid-phase catalyst essential to obtain a gel-free product. Soluble 25 catalysts, such as, for example, the biscyclopentadienyltitanium(IV) dichloride-aluminum alkyl system, are generally unsuitable because the second terminal bond of the  $\alpha, \omega$ -diene is not protected from the catalyst and remains available for reaction in a 30 gel-forming polymerization reaction. Heterogeneous catalysts capable of producing stereo-regular polymers are also preferred, and particularly those catalysts suitable for the preparation of insoluble, isotactic polypropylene. Titanium catalyst containing a minor 35 amount of prepolymerized α-olefin and solid-supported titanium catalysts particularly are preferred. Prepolymerized activated titanium catalyst is described

in U. S. Patent 4,295,991 to Wristers which is hereby incorporated herein by reference. Briefly, the prepolymerized activated titanium catalyst is obtained by reducing titanium trichloride with an organoaluminum compound at low temperatures, treating the resulting reduced solid product with about 1-1000 weight percent α-olefin, and activating the resulting prepolymerized solid with a halogenated hydrocarbon and a Lewis base complexing agent or titanium tetrachloride and a Lewis base complexing agent. The prepolymerized activated titanium catalyst may be employed in combination with an organometal compound.

Specific examples of supported titanium catalysts include: (1) a catalytic solid powder obtained by co-15 pulverizing a magnesium compound such as magnesium chloride in an electron donor such as aromatic carboxylic esters or by reacting these compounds optionally in the presence of an inert solvent, and then co-pulverizing the resulting product together with 20 a titanium compound; (2) a pulverulent solid obtained by reacting a reaction product of a titanium compound and an electron donor with a magnesium compound by co-(3) a pulverulent solid pulverizing the compounds; obtained by reacting a magnesium compound, an electron 25 donor and a titanium compound together through the simultaneous co-pulverization of these compounds; a product obtained by supporting titanium tetrachloride on a co-pulverized product of a solid reaction product [(hereinafter referred to as a support magnesium hydroxychloride and ethylaluminum dichloride and an aromatic carboxylic acid ester; (5) a product obtained by co-pulverizing the support (I) and a complex of an aromatic carboxylic acid ester and (6) a product obtained by titanium tetrachloride; 35 treating the co-pulverized product obtained in (5) in a halogenated hydrocarbon with halide such as halogen, interhalogen compound, sulfur monochloride, sulfur

dichloride or phosphorous pentachloride; (7) a product obtained by treating a copulverized product of the support (I) and an aromatic acyl halide with titanium tetrachloride and ethers or an alkoxy compound; (8) a 5 product obtained by treating a copulverized product of the support (I) and an alkoxysilane with titanium tetrachloride and phosphates or phosphites; product obtained by subjecting a solid product [a support (II)] obtained by reacting the support (I) with 10 a siloxane compound, instead of the support (I), to any procedure described in (4) through (8); (10) a product obtained by using substantially anhydrous magnesium chloride [a support (III)], instead of the support (I), in any procedure described in (4) through (8); (11) a 15 product obtained by adding a halide such as silicon tetrachloride. trichloride, aluminum phosphorus pentachloride, antimony trichloride and tellurium tetrachloride, for example, during the co-pulverization in any procedure described in (4) through (8); (12) a 20 product obtained by adding a siloxyaluminum compound such as trimethylsiloxyaluminumdichloride during the co-pulverization in any procedure described (4) through (10); (13) a product obtained by adding siloxane compound during the co-pulverization in any 25 procedure described in (4) through (10); (14) a solid obtained by reacting the reaction product of a Grignard compound and a compound containing a Si-H bond, alcohols or phenols, siloxanes, with a containing a Si-halogen bond, reacting the resulting 30 solid [a support (IV)] with titanium tetrachloride, and finally reacting the resulting product with carboxylic acid ester; (15) a solid powder obtained by treating a reaction product of magnesium oxychloride and ethylaluminumdichloride with a polysiloxane, and 35 reacting the resulting product with an alcohol and diethylaluminumchloride in the order described, and finally reacting the resulting product with titanium

(16) a solid powder obtained by tetrachloride; reacting magnesium chloride pulverized in a ball mill with an alkyl titanate and then reacting the resulting product with titanium tetrachloride and methyl hydrogen 5 polysiloxane; (17) a solid powder obtained by treating a co-pulverized product of magnesium chloride and ethoxytrichlorotitanium with iodine trichloride in a halogenated hydrocarbon; (18) a solid powder obtained by using magnesium chloride in the procedure described 10 in (15) instead of a compound obtained by treating a product of magnesium oxychloride reaction ethylaluminum dichloride with a polysiloxane; solid powder obtained by using a reaction product of magnesium oxychloride and ethylaluminum dichloride in 15 the procedure described in (16), instead of magnesium Additional examples of solidchloride; and the like. supported catalysts are described in U. S. Patent 4,366,296 which is hereby incorporated herein by reference.

As mentioned above, the selection of catalyst can used to favor the formation of side chain unsaturation or long chain branching. Generally, the prepolymerized catalyst has been found to favor more side chain unsaturation, while the magnesium chloride-25 supported catalysts tend to favor more long chain branching.

The copolymerization is effected in gas or liquid phase, and in the case of propylene, preferably as a The copolymerization slurry in liquid propylene. 30 should be in the essential absence of a solvent, which commonly include pentane, hexane, heptane, octane, branched alkanes such as those sold under the trade Such solvents tend to designation Isopar or the like. adversely affect the copolymer by permitting cross-35 linking and gel formation to occur which is believed to result from the solubility of the copolymer therein and the ability of the side-chain unsaturation to WO 91/17194 PCT/US91/03141

participate in a dissolved state in subsequent crosslinking polymerizations. Nonetheless, a very minor proportion of solvent, or a diluent by which the copolymer is not swollen, may permissibly be present in the reactor, e.g. the catalyst may be introduced to the reactor in such a solvent, without significant adverse effects.

The copolymerization reactor temperature must be maintained below the crystalline melting point of the propylene copolymer to facilitate inhibiting the mobility of the unsaturated side chain in participation in additional polymerization reactions which cross-link the polymer to form xylene-insoluble gel. The crystalline melting point of the copolymer is usually in the range of about 155-165°C. The reactor is preferably maintained at from 0° to 100°C, and especially from 30° to 70°C.

In preparing the copolymer, the propylene and a, wdiene are charged to a suitable reactor vessel with the
catalyst. Generally, the propylene in gas or liquid
form comprises the bulk of the reactor contents and the
diene comonomer(s) may comprise from about 0.001 to
about 20 weight percent. If the proportion of diene
comonomer feed is too low, insufficient comonomer is
incorporated in the resultant copolymer. With an
excessive diene comonomer proportion, however, the
comonomer can begin to function as a diluent or solvent
for the copolymer and gel formation can result.
Preferably, the reactor charge comprises from about
0.01 to about 20 weight percent of the diene comonomer,
especially from about 0.1 to about 10 weight percent.

The reactor charge may also include hydrogen for controlling molecular weight and improving catalyst activity. The hydrogen concentration may be as high as 10 mole percent, preferably from about 0.01 to about 5 mole percent, and especially from 1 to 2 mole percent. In general, the more hydrogen present in the reaction

zone, the lower the molecular weight and the greater the catalyst activity. Catalyst is generally fed to the reactor in suitable amounts to obtain the desired copolymer production. The reactor may be operated continuously, batch or semi-batch, at a suitable pressure, usually from about 50 to about 2000 psig, preferably from about 100 to about 700 psig, and more preferably from about 300 to about 600 psig.

It is contemplated that the copolymer of the present invention may be used as a reactive feed for subsequent polymerization reactors, such as, for example, the ethylene-propylene rubber reactor in a polypropylene impact copolymer process.

The copolymer is recovered from the reactor and 15 separated from unreacted monomer in a conventional manner. Because of its novel structure, a wide variety of end-use applications are contemplated. chain unsaturation imparts stability to the copolymer not possessed by homopolypropylene. For example, the 20 propylene-diene copolymer suffers no molecular weight loss (increased melt flow rate) and forms gel when treated with peroxide free radical initiators, or when exposed to gamma radiation, whereas conventional polypropylene rapidly degrades through a molecular 25 chain scission mechanism and does not form gel. the present copolymer may be grafted with vinyl monomers such as maleic anhydride using peroxide initiators, or exposed to gamma radiation. This later application is particularly attractive in medical 30 devices and structures made from or containing the present copolymer, e.g. syringes, trays and the like, wherein gamma radiation, typically 2.5-10 MRads, is for sterilization purposes. Indeed, radiation serves to cross-link the present propylene 35 copolymer via the unsaturation in the side chains, conferring additional dimensional stability on the medical device or other product.

The α,ω-diene-propylene copolymer also enhanced processability owing to the presence of the long chain branching, in much the same manner that branched polyethylene is known to be easier to process than linear polyethylenes. For example, in food packaging applications wherein the polypropylene container is retorted, e.g. a bottle or dish, the propylene-diene copolymer is easily blow molded or stretch blow molded, whereas conventional polypropylene has such poor melt strength that thin, non-uniform 110 walls are usually obtained in the molded products.

In pipe fabrication, the present propylene-diene copolymers have better processability then conventional polypropylene of equivalent melt flow rate. 15. copolymer has a lower melt viscosity at a given shear rate because of the long side chains, allowing higher throughput and less sag at the extruder die. Also, the copolymer can be extrusion coated with less neck-in than conventional polypropylene, avoiding the necessity of blending with low density polyethylene as 20 conventionally done with the propylene homopolymer, and avoiding the concomitantly reduced maximum temperatures. Moreover, the a, w-diene-propylene copolymers retain flexural modulus better than conventional polypropylene, allowing the use of the 25 copolymer at higher temperatures in applications where the prior art polypropylene would be unsatisfactory.

#### EXAMPLES

30 In the examples that follow, the catalysts are designated as follows: "catalyst I" is prepolymerized TiCl<sub>3</sub> catalyst prepared in accordance with description in Example 10 of U. S. Patent 4,295,991; and "catalyst II" is prepolymerized MgCl2-supported 35 alkoxysilane-modified TiCl catalyst triethylaluminum as a cocatalyst, wherein the catalyst

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is obtained from Mutsui Petrochemical Company under the trade designation SP-101. 1,9-Decadiene from Shell International Chemicals was purified by passage over a column of basic alumina (Baker, Brockman activity 5).

The column was 1 inch in diameter and 10 inches high. The diene was collected over activated 4 Å molecular sieves and sparged with nitrogen for 1 hour in order to remove air and water.

1,9-Decadiene contents of the propylene copolymers

were measured by infrared spectroscopy. One gram of
the copolymer sample was dissolved in 100 ml of xylene.
The copolymer-xylene solution was then poured into 500
ml of cold isopropanol. The resulting slurry was
filtered and the solid dried under vacuum for 16 hours
at 70°C. A 10 mil pad was compression molded at 190°C
and this plaque was used for the infrared measurements.
Unsaturation in the side chains was determined by
measuring the absorbance at 1640 cm-1 and the sample
thickness and applying the equation:

# $x_{unsat} = k\lambda_{1640}/\delta$

wherein  $x_{usset}$  is the weight percent of unsaturated diene comonomer on the copolymer, k is a constant,  $A_{1640}$  is absorbance at 1640 cm<sup>-1</sup>, and  $\delta$  is the sample thickness in mm. The total amount of 1,9-decadiene incorporated into the copolymer was also determined by infrared spectroscopy. This was calculated by measuring the absorbance at 722 cm<sup>-1</sup> and the sample thickness and applying the equation:

# $x_{diene} = kA_{722}/\delta$

wherein  $x_{\text{diene}}$  is the weight percent of diene comonomer in the copolymer,  $A_{722}$  is the absorbance at 722 cm<sup>-1</sup> and k and  $\delta$  are as above. The amount of 1,9-decadiene bridging between two polymer chains was determined by

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subtracting the amount with terminal unsaturation from the total 1,9-decadiene content. The unsaturation content of the polymer was also determined in some cases by high temperature proton nuclear magnetic resonance spectroscopy of the sample dissolved in tetrachloroethane. The mole percentage unsaturation was calculated using the following equation:

# $y_{unsat} = 100 [2A/(B-3A)]$

wherein yunsat is the unsaturated diene mole percentage of the sample, A is the integral of the spectrum from 5 10 to 6 ppm (olefinic region) and B is the integral of the spectrum from 0 to 2 ppm (aliphatic region).

The average molecular weight of the polymers was determined by gel permeation chromatography (GPC). number of long side chains/molecule for a given weight average molecular weight was then calculated from the number of bridges and the weight average molecular weight.

The melt flow rate (MFR) of the sample was determined by ASTM D-1238, condition L (2.16 kg, 230°C). The heptane insolubles (HI) were determined by measuring the fraction of polymer which remained in an extraction thimble following extraction with refluxing heptane in a Soxhlet extractor under nitrogen for 24 The melting point (MP) was determined by 25 hours. differential scanning calorimetry, based on a second melt. Both the heating and cooling rates were 10°C/min.

#### COPOLYMER PREPARATION AND PROPERTIES

# 30 Example 1

A dry, nitrogen-blanketed two-liter autoclave was charged with 100 ml of 1,9-decadiene; 4 ml of 1 M triethylaluminum; 4 ml of 0.1 M alkoxy silane catalyst modifier; hydrogen to 110 psig; and 1250 ml of liquid

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propylene. The autoclave was continuously stirred and heated to 60°C. Two hundred mg of catalyst I was pressured into the autoclave and polymerization allowed to proceed at 70°C for 1.5 hours. The autoclave was then vented and cooled, and a granular polymer was The product was placed in a vacuum oven recovered. (50°C, 20 mm Hg) overnight. The dried polymer product weighed 261 g. Proton nuclear magnetic resonance spectrometry showed 0.55 mole percent unsaturation, in contrast to polypropylene similarly prepared which contains no measurable unsaturation.

# Example 2

Propylene/1,9-decadiene copolymers were produced in two 150 gallon stirred tanks, operated in series, as a reactor. Catalyst I was used. The conditions in the two reactors were as follows:

		REACTOR 1	REACTOR 2
	Reaction Temperature ('F)	169.8	159.3
20	Propylene Flow Rate (lb/hr) 1,9-Decadiene Flow Rate	109.9	50.0
	(lb/hr) Gas Phase H <sub>2</sub> Concentration	5.3	2.4
	<pre>(mole %)  Solids Concentration (wt %)</pre>	3.0 29.9	3.3 30.2

25 Sufficient catalyst was fed to produce polymer at a rate of 50.6 lb/hr. Sufficient 15% diethylaluminum chloride (DEAC) in hexane was fed to maintain an Al/Ti ratio of 6.0. The product had a MFR of 27.5 dg/min and an HI of 87.5%. It contained 4.4 ppm Ti. copolymer had a melting point of 156.6°C and contained 30 0.30 mole percent 1,9 decadiene unsaturation and 0.50 mole percent total 1,9-decadiene. GPC measurement of the molecular weight distribution in this product gave an  $M_w = 186,700$  and  $M_w/M_n = 5.26$ . The measured 35 intrinsic viscosity of the sample was 0.833. intrinsic viscosity calculated from the GPC data was This gave a branching index of 0.674. 1.235.

demonstrates that significant branching was present in the copolymer granules from the polymerization reactor. However, the sample contained no gel and was completely soluble in refluxing xylene. The effect of branching evident was also from recoverable compliance measurements of the polymer. The recoverable compliance at 180°C was 6.26 x 10<sup>-5</sup> cm<sup>2</sup>/dyne. of the polymer were injection molded into 125 mil thick bars and subjected to gamma radiation at 1 and 5 MRad dosages. Before irradiation, the gel content was 0%. After irradiation at 1 MRad and 5 MRads, the gel contents were 3% and 34%, respectively. The MFR was also measured on ground injection molded bars, before and after irradiation. Before irradiation, the MFR was 38 dg/min. After irradiation at 1 MRad and 5 MRads, the MFR's were 31 and 2.6 dg/min, respectively. experiments show that the molecular weight increased due to cross-linking of the polymer after exposure to radiation, in contrast to propylene homopolymers which exhibit severe molecular weight loss with such gamma radiation exposures.

#### Example 3

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Propylene/1,9-decadiene copolymers were produced in the same reactor as described in Example 2. Catalyst system I was used. The conditions in the two reactors were as follows:

		REACTOR 1	REACTOR 2
	Reaction Temperature (°F)	169.8	160.2
30	Propylene Flow Rate (lb/hr) 1,9-Decadiene Flow Rate	110.0	50.0
	(lb/hr) Gas Phase H <sub>2</sub> Concentration	7.9	3.6
	(mole %) ~	1.5	1.7
	Solids Concentration (wt %)	33.6	33.1

Sufficient catalyst was fed to produce polymer at a rate of 56.7 lb/hr. Sufficient 15% DEAC in hexane was fed to maintain an Al/Ti ratio of 4.7. The product had a MFR of 4.2 dg/min and an HI of 90.0%. It contained 5 . 7.2 ppm Ti. The polymer had a melting point of 157.9°C and contained 0.10 mole % 1,9-decadiene unsaturation and 0.50 mole percent total 1,9-decadiene. measurement of the molecular weight distribution in this product gave a weight average molecular weight (M\_) of 250,000 and ratio of number average molecular 10 weight (Mn) to M, of 5.40. The intrinsic viscosity of sample was 1.114. The intrinsic viscosity calculated from the GPC data was 1.580. This gave a branching index of 0.705. Significant branching was present in the polymer granules from the polymerization 15 reactor. The gel content of the sample was 0.0%. recoverable compliance at 180°C was 6.56 x cm<sup>2</sup>/dyne. Samples of the polymer were injection molded into 125 mil thick bars and subjected to gamma 20 radiation. Before irradiation, the gel content was 0%. After irradiation at 1 MRad and 5 MRads, the gel contents were 19% and 43%, respectively. The MFR was measured on ground injection molded bars, before and after irradiation. Before irradiation, the MFR was 8 After irradiation at 1 MRad, the MFR was 2 dq/min. 25 After exposure to 5 MRads of radiation, the dg/min. sample MFR could not be measured since it would not flow through an orifice. These experiments show that the molecular weight increased due to cross-linking of 30 the polymer after exposure to radiation.

## Example 4

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Propylene/1,9-decadiene copolymers were produced in the same reactor as described in Example 2. Catalyst system I was used. The conditions in the two reactors were as follows:

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		REACTOR 1	REACTOR 2
	Reaction Temperature ('F) Propylene Flow Rate (lb/hr) 1,9-Decadiene Flow Rate	169.8 110.0	159.4 50.0
5	(lb/hr) Gas Phase H <sub>2</sub> Concentration	7.2	3.3
	<pre>(mole %) T Solids Concentration (wt %)</pre>	1.8 30.6	1.9

Sufficient catalyst was fed to produce polymer at a rate of 56.9 lb/hr. Sufficient 15% DEAC in hexane was fed to maintain an Al/Ti ratio of 4.7. The product had a MFR of 7.0 dg/min and an HI of 89.0%. It contained 6.8 ppm Ti. The polymer had a melting point of 158.1°C and contained 0.10 mole percent 1,9-decadiene unsaturation and 0.50 mole percent total 1,9-decadiene.

# Comparative Example 1

A commercial propylene homopolymer with an MFR of 1.6 dg/min was irradiated at different dosages to demonstrate the degradation that results when propylene homopolymers are subjected to gamma radiation. samples were injection molded into 125 mil Izod impact bars and subjected to gamma radiation. After 2.1 MRads the MFR increased to 8 dg/min, at 5.6 MRads the MFR increased to 20 dg/min, and after 10.6 MRads the MFR 25 was 78 dg/min. No cross-linking was observed.

# Comparative Example 2

A propylene/hexene copolymer with an MFR of 2.5 dg/min irradiated at different dosages to demonstrate that severe degradation results 30 propylene copolymers with other  $\alpha$ -olefins are subject to gamma radiation. After 2.1 MRads the MFR increased to 10 dg/min. and at 5.6 MRads the MFR increased to 39 dg/min.

# Comparative Example 3

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A propylene homopolymer with a MFR of 15 dg/min was irradiated at different dosages to demonstrate the degradation that results when propylene homopolymers are subjected to gamma radiation. The samples were injection molded into 125 mil Izod impact bars and subjected to gamma radiation. Before irradiation the gel content of the sample was 0.0%. After irradiation at 2.1 MRads, 5.6 MRads, and 10.6 MRads the gel content was 0.0%. No cross-linking was observed.

# Comparative Example 4

This example demonstrates the low recoverable compliance that is observed for polypropylene homopolymers. The recoverable compliance homopolymer having a MFR of 8 dg/min was measured at The recoverable compliance was  $2.80 \times 10^{-5}$ This demonstrates the lower elasticity of cm<sup>2</sup>/dyne. the melt for a linear homopolymer.

The foregoing description of the invention is illustrative and explanatory only. Various modifications will occur to the skilled artisan in view thereof. All such variations which are within the scope or spirit of the appended claims are intended to be embraced thereby.

## Claims:

- 1. An α-olefin/α,ω-diene copolymer, comprising: an α-olefin having from 2 to about 12 carbon atoms, interpolymerized with from 0.01 to 5 mole percent of an α,ω-diene having from 7 to about 30 carbon atoms, to produce an unsaturated crystalline copolymer essentially free of gel.
- 2. The copolymer of claim 1, wherein the  $\alpha$ -olefin comprises propylene.
- 3. The copolymer of claim 1, comprising from about 0.05 to about 2 mole percent of said  $\alpha, \omega$ -diene.
- 4. The copolymer of claim 1, comprising from abut 0.001 to about 5 mole percent of unsaturated  $\alpha, \omega$ -diene.
- 5. The copolymer of claim 1, wherein the  $\alpha, \omega$ -diene has from 8 to 12 carbon atoms.
- 6. An unsaturated propylene/ $\alpha$ ,  $\omega$ -diene copolymer, comprising:

propylene interpolymerized with from 0.01 to 5 mole percent of an  $\alpha, \omega$ -diene having from 8 to 12 carbon atoms, wherein the copolymer contains unsaturated side chains, is essentially crystalline and is essentially free of gel.

- 7. The copolymer of claim 6, wherein said copolymer comprises from about 0.05 to about 2 mole percent of said  $\alpha, \omega$ -diene.
- 8. The copolymer of claim 6, wherein said copolymer comprises from about 0.001 to about 5 mole percent unsaturated side chains.
- 9. The copolymer of claim 6, wherein copolymer further includes a minor proportion of ethylene.
- 10. The copolymer of claim 6, wherein the diene comprises 1,9-decadiene.

- 11. The composition of claim 6, wherein said copolymer has a xylene-insoluble fraction of not more than 0.1 weight percent.
- 12. The composition of claim 6, wherein said copolymer has a heptane-insoluble fraction of at least 80 weight percent.
- 13. The composition of claim 6, wherein said copolymer has a branching index of from about 0.2 to about 0.9.
- The copolymer of claim 6, wherein the copolymer is 14. prepared by polymerization with heterogeneous catalyst for stereoregular polymerization at a temperature below crystalline melting point of the copolymer essentially in the absence of a solvent.
- 15. A propylene/decadiene copolymer, comprising:
  - propylene copolymerized with from 0.01 to 5 mole percent total 1,9-decadiene and from about 0.001 to about 5 mole percent unsaturated 1,9-decadiene wherein the copolymer has less than 0.1 weight percent of xylene-insolubles, at least 80 weight percent heptane insolubles and a branching index of about 0.2 to about 0.9.
- 16. The copolymer of claim 15, comprising from 0.01 to 2 mole percent total 1,9-decadiene.
- 17. The copolymer of claim 15, comprising from 0.01 to 2 mole percent unsaturated diene.
- 18. The copolymer of claim 15, comprising from 80 to 95 weight percent heptane insolubles.
- 19. The copolymer of claim 15, wherein the copolymer has a branching index of from about 0.6 to about 0.8.

20. A method for preparing a gel-free unsaturated copolymer of an  $\alpha$ -olefin and an  $\alpha, \omega$ -diene, comprising:

charging to a reaction zone an  $\alpha$ -olefin having from 2 to about 12 carbon atoms, from 0.01 to 20 weight percent of an  $\alpha, \omega$ -diene having from 7 to about 30 carbon atoms, and a solid heterogeneous coordination catalyst;

maintaining the reaction zone essentially free of solvent and at atemperature at which the α-olefin is fluid but below the crystalline melting point of the copolymer, effective to produce an essentially gel-free, linear copolymer with unsaturation.

- 21. The method of claim 20, wherein the α-olefin is selected from ethylene, propylene, butene-1, pentene-1 and 4-methyl-1-pentene.
- 22. The method of claim 20, wherein the  $\alpha,\omega$ -diene has up to 12 carbon atoms.
- 23. The method of claim 20, wherein the catalyst is a Ziegler-type catalyst.
- 24. The method of claim 20, wherein up to 10 mole percent hydrogen is charged to said reaction zone to control molecular weight of said copolymer.
- 25. The method of claim 20, wherein the copolymer has from 0.1 to 1 mole percent unsaturation.
- 26. A method for preparing a gel-free, unsaturated copolymer of an  $\alpha$ -olefin and an  $\alpha, \omega$ -diene, comprising:

charging an  $\alpha$ -olefin selected from ethylene, propylene, butene-1,pentene-1 and 4-methyl-1-pentene to a reaction zone with from 0.01 to 20 weight percent of an  $\alpha,\omega$ -diene of from 7 to 12 carbon atoms, up to 10 mole percent hydrogen, and a

solid, heterogeneous Ziegler-type catalyst;

- maintaining the reaction zone essentially free of solvent and at atemperature at which the  $\alpha$ -olefin is fluid but substantially below the crystalline melting point of the copolymer; and
- recovering an essentially linear and gel-free copolymer from the reaction zone, wherein the copolymer has from 0.1 to 1 mole percent unsaturation.
- 27. The method of claim 26, wherein the  $\alpha$ -olefin is propylene.
- 28. The method of claim 26, wherein the  $\alpha, \omega$ -diene comprises 1,9-decadiene.
- 29. The method of claim 26, wherein the catalyst is a solid-supported catalyst.
- The method of claim 26, wherein the catalyst is 30. obtained by (1) contacting TiCl, with organoaluminum compound at a temperature between about -50' and 30'C to produce a TiCl, reduced solid product, (2) contacting the reduced solid TiCl, product with a minor amount of an a-olefin having at least 3 carbon atoms polymerization conditions to obtain a reduced TiCl, solid product containing between about 1 and 1000 weight percent of prepolymerized α-olefin based on the weight of TiCl3, and (3) treating said prepolymerized reduced solid with one of (a) a chlorinated hydrocarbon having at least 2 carbon atoms and a Lewis base complexing agent or (b) TiCl, and a Lewis base complexing agent, to convert the prepolymerized TiCl, reduced solid to substantially non-friable highly crystalline prepolymerized TiCl, composition.
- 31. The method of claim 30, wherein the catalyst further comprises an organometallic cocatalyst.

32. A method of preparing an isotactic, essentially gel-free copolymer of propylene and an  $\alpha,\omega$ -diene comprising:

charging propylene, from 0.01 to 20 weight percent 1,9-decadiene, from 0.01 to 5 mole percent hydrogen, and a solid heterogeneous Ziegler-type catalyst for isotactic polymerization, to a reaction zone;

maintaining the reaction zone essentially free of solvent and at a temperature below about 150°C effective to randomly copolymerize said propylene and said diene; and

recovering an essentially gel-free isotactic copolymer.

- 33. The method of claim 32, wherein a minor proportion of ethylene is charged to said reaction zone.
- 34. The method of claim 32, further comprising polymerizing said copolymer with ethylene and propylene in another rection zone.
- The method of claim 32, wherein said catalyst is 35. obtained by (1) contacting TiCl with organoaluminum compound at a temperature between about -50° and 30°C to produce a TiCl3 reduced solid product, (2) contacting the reduced solid  $TiCl_3$  product with a minor amount of an  $\alpha$ -olefin having at least carbon atoms under polymerization conditions to obtain a reduced TiCl3 solid product containing between about 1 and 1000 weight percent of prepolymerized  $\alpha$ -olefin based on the weight of TiCl3, and (3) treating said prepolymerized reduced solid with one of (a) a chlorinated hydrocarbon having at least 2 carbon atoms and a Lewis base complexing agent or (b) TiCl, and a Lewis base complexing agent, to convert the prepolymerized TiCl; reduced solid to

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- a substantially non-friable highly active, crystalline prepolymerized TiCl<sub>3</sub> composition.
- 36. The method of claim 35, wherein the catalyst includes an organometallic cocatalyst.

# INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 91/03141

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According to Int. Cl	to International Patent l . 5	Classification (IPC) or to both Nationa C 08 F 210/00	l Classification and IPC	•
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Category °	Citation of Do	cument, 11 with indication, where approp	priate, of the relevant passages 12	Relevant to Claim No.13
x	S S J I	HEMICAL ABSTRACTS, vo eptember 1969, (Colum ee page 23, abstract P-A-69 06 275 (MITSUI NDUSTRY CO., LTD) 17 I he abstract	bus, Ohio, US), no. 39638e, & CHEMICAL	1-9
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# ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

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This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 10/09/91

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